PATENT COOPERATION TREATY

REC'D 2 7 JUN 2005

NTERNAT	IONAL SEARCH	IING AUTH	ORITY		WIPO	PC
To: LAURA A. CORUZZI JONES DAY					PCT	<u>, , , , , , , , , , , , , , , , , , , </u>
222 EAST	41ST STREET			WR	ITTEN OPINION OF THE	
NEW YORK, NY 10017-6702				INTERNATIO	ONAL SEARCHING AUTHORITY	
					(PCT Rule 43bis.1)	
				Date of mailing (day/month/year)	22 JUN 2005 .	
	's or agent's file re	oforence		FOR FURTHER	ACTION See paragraph 2 below	
10589-13-	228 nal application No		International filing date	(day/month/vear)	Priority date (day/month/year)	一
	••				27 March 2003 (27.03.2003)	
PCT/US0	1/09572 nal Patent Classifi	cation (IPC)	26 March 2004 (26.03.2 or both national classificat		27 March 2003 (27.03.2003)	ᅥ
	01N 61/00; C12Q				7.21, 41, 69.2, 91.3, 183; 514/1, 2	_
	RAPEUTICS, IN	D				
1. This	opinion contains i	ndications rel	ating to the following item	as:		
\boxtimes	Box No. I	Basis of the	opinion			
Box No. II Priority						
Box No. III Non-establishment of opinion with regard to novelty, inventive step and industrial applicability			ative step and industrial applicability			
Box No. IV Luck of unity of invention						
Box No. II Non-establishment of opinion with regard to novelty, inventive step and industrial applicability Box No. IV Lack of unity of invention Box No. V Reasoned statement under Rule 436i:1(a)(i) with regard to novelty, inventive step or industrial applicability, citations and explanations supporting such statement				novelty, inventive step or industrial atoment		
	Box No. VI	Certain do	cuments cited			1
	Box No. VII	Certain del	fects in the international ap	plication		
Box No. VIII Certain observations on the internation			servations on the internation	onal application		
	THER ACTIO					
Inter Auth	national Prelimin ority other than t	ary Examini nis one to be	ng Anthority ("IPEA") e	xcept that this does IPEA has notified th	be considered to be a written opinion of the not apply where the applicant chooses an se International Bureau under Rule 66.1bis(b) ered.	ı l
If this opinion is, as provided above, considered to be a written opinion of the IPEA, the applicant is invited to submit to the IPEA a written reply together, where appropriate, with amendments, before the expiration of 3 months from the date of mailing of Form FUTSA/220 or before the expiration of 22 months from the pricrity date, whichever expires later.						
For f	For further options, see Form PCT/ISA/220.					
3. For f	urther details, sec	notes to For	m PCT/ISA/220.			
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	l mailing address Mail Stop PCT, Att		0	Valer	ie Bell-Harrisi	2

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Mail Stop FCT, Attn. ISA/US
Commissioner for Patents
P.O. Box 1450
Alexandrie, Virginia 22313-1450
Facsimile No. (703) 305-3230
Form PCT/ISA/237 (cover sheet) (January 2004)

International application No.
PCT/US04/09572

Box No	o. I Basis of this opinion				
With regard to the language, this opinion has been established on the basis of the international application in the language in which it was filed, unless otherwise indicated under this item.					
	This opinion has been established on the basis of a translation from the or which is the language of a translation furnished for the purposes of intern	riginal language into the following language, ational search (under Rules 12.3 and 23.1(b)).			
2. With inven	regard to any nucleotide and/or amino acid sequence disclosed in the in tion, this opinion has been established on the basis of:	ternational application and necessary to the claimed			
a.	type of material	·			
	a sequence listing				
	table(s) related to the sequence listing				
ь.	format of material				
	in written format				
	in computer readable form				
c.	time of filling/furnishing				
	contained in international application as filed.				
	filed together with the international application in computer reada	ble form.			
	furnished subsequently to this Authority for the purposes of search	<u>.</u>			
3. 🗌	In addition, in the case that more than one version or copy of a sequent or furnished, the required statements that the information in the subsecupilication as filed or does not go beyond the application as filed, as app	quent or additional copies is identical to that in the			
4. Addit	tional comments:				
	·				
	·				

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Box No. III Non-establishment of opinion with regard to novelty, inventive step and industrial applicability

The questions whether the claimed invention appears to be novel, to involve an inventive step (to be non-obvious), or to be industrially applicable have not been examined in respect of:				
	the entire international application	l		
\boxtimes	claims Nos. <u>35 and 52</u>	-		
becar	use:			
	the said international application, or the said claim Nos relate to the following subject matter which does not require an international preliminary examination (specific):			
\boxtimes	the description, claims or drawings (indicate particular elements below) or said claims Nos. 35 and 52 are so unclear that no meaningful opinion could be formed (specify). Claims 35 and 52 are multiple dependent claims that depend from claims 33 and 34, which are dependent from claim 12, which is a multiple dependent claim. Thus a multiple dependent claim (i.e., claim 12) serves as a basis for claims 35 and 52, which are multiple dependent claims. Claims 35 and 52, which are multiple dependent claims. Claims 35 and 52, threshre, are improper dependent claims, (see Rule 6.4 (a)).			
	the claims, or said claims Nos are so inadequately supported by the description that no meaningful opinion could be formed. In international search report has been established for said claims Nos the nucleotide and/or amino acid sequence listing does not comply with the standard provided for in Annex C of the Administrative Instructions in that: the written form has not been furnished does not comply with the standard the computer readable form has not been furnished			
	does not comply with the standard the tables related to the nucleotide and/or amino acid sequence listing, if in computer readable form only, do not comply with the technical requirements provided for in Amex C-bie of the Administrative Instructions. See Supplemental Box for further details.			

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Box No. IV Lack of unity of invention				
In response to the invitation (Form PCT/ISA/206) to pay additional fees the applicant has: paid additional fees				
paid additional fees under protest				
not paid additional fees				
 This Authority found that the requirement of unity of invention is not complied with and chose not to invite the ap pay additional fees. 	plicant to			
 This Authority considers that the requirement of unity of invention in accordance with Rule 13.1, 13.2 and 13.3 is 				
complied with				
not complied with for the following reasons:				
See the lack of unity section of the International Search Report(Form PCT/ISA/210)	-			
•				
4. Consequently, this opinion has been established in respect of the following parts of the international application: all parts.				
the parts relating to claims Nos. <u>1-34,36-51,53 and 54</u>				

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	Box No. V Reasoned statement under Rule 43 bis.1(a)(i) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement			
1	1. Statement			
	Novelty (N)	Claims 1-32 and 40-51	YES	
		Claims 33, 34, 36-39, 53 and 54	NO	
	Inventive step (IS)	Claims NONE	YES	
		Claims 1-34, 36-51, 53, 54	NO	
	Industrial applicability (IA)	Claims 1-34, 36-51, 53, 54	YES	
		Claims NONE	NO	

2. Citations and explanations:

Please See Continuation Sheet

Claims 1-34, 36-51, 53 and 54 meet the criteria set out in PCT Article 33(4), and thus possess industrial applicability because the subject matter claimed can be made or used in industry.

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Box No. VIII Certain observations on the international application

The following observations on the clarity of the claims,	description, and drawings or on the questions whether the claims are full	ly
supported by the description, are made:		

Claims 35 and 52 are multiple dependent claims that depend from claims 33 and 34, which are dependent from claim 12, which is a multiple dependent claim. Thus a multiple dependent claim at 25, which are multiple dependent claims. Claims 35 and 52, which are multiple dependent claims. Claims 35 and 52, therefore, are improper dependent claims, (see Rule 6.4 (a)).

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In case the space in any of the preceding boxes is not sufficient.

V. 2. Citations and Explanations:

Claims 33, 34, 36-39, 53 and 54 lack novelty under PCT Article 33(2) as being anticipated by US 6,446,032 B1 (SCHIMMEL).

Schimmel discloses small molecule, (e.g., see bottom of col. 27-28), antiproliferative, (e.g., chemothempeutic agents: see col. 3, compounds for trenting cancer when administered to a bost (e.g., burnou). These RNA (e.g., RNA) binding compounds compress structure within the scope of the presently claimed invention (e.g., see col. 27-28, examples and patent claims). The ability to inhibit RNA aplicing compoundses such services to the ability of these compounds to bind RNA. A nay event, the claims is not structure-limited and the PTO locks the facilities for making comparisons between prior art compounds and the claimed prospective assay-derived compounds.

Claims 33, 34, 36-39, 53 and 54 lack novelty under PCT Article 33(2) as being anticipated by WO 01/25486 A1 (RANA).

Rana discloses assay-derived RNA inhibiting (e.g., binding; see e.g., bottom of page 9400 of page 10; and claims, especially claims 1; 2, 28-30, 04-30 compounds within the scope of the presently claimed invertine (e.g., claims 2-52) that are antipromilibrative for use in treating proliferative disorders (e.g., cancer; i.e., see claim 46) when administered to humans. The ability to his distribution of the properties of the present due to the ability of these compounds to bind RNA (e.g. RNA). In any event, the claim is not structure-limited and the PTO lacks the facilities for making comparisons between prior art compounds and the claimed and the claimed prospective assets—drived commons.

Claims 33, 34, 36-39, 53 and 54 lack novelty under PCT Article 33(2) as being anticipated by WO 02/083837 A1 (ALMSTEAD).

Almstead discloses assay-durived binding compounds (e.g. see pages 3-4, bottom of page 10-11) within the scope of the presently claimed invention (e.g. see pages 21-25, daimed just any time autiprolibrative for see in treating prolibrative disorders (e.g., cancer) when administered to humans. The ability to inhibit RNA spiking endomulcless is inherently present due to the ability of these compounds to bind RNA (e.g. RNA). In any event, the claim is not structure-limited and the PTO lacks the fieldlities for making comparisons between prior art compounds and the claimed prospective assocyacierved compounds.

Claims 33, 34, 36-39, 53 and 54 lack novelty under PCT Article 33(2) as being anticipated by WO 02/083953 A1 (RANDO et al.).

Rando et al. disclose assay-derived RivA binding (e.g., RNA) compounds which effect RNA bost cell factor complexes in vivo.

(e.g. RNA aplicing see page 10; bottom of page 12-page 13) which compounds are within the scope of the presently claimed invention
(e.g. see claim 3) that are antiproliferative for use in treating proliferative disorders (e.g., cancer) when administered to humans. The
shifty to inhibit RNA applicing endouncleases is inherently present due to the ability of these compounds to bind RNA (e.g. RNA). In
any event, the claim is not structure-limited and the PTO locks the facilities for making comparisons between prior art compounds and
the claimed prospective assay-derived compounds.

Claims 1-34, 36-51, 53 and 54 lack an inventive step under PCT Article 33(3) as being obvious over WO 01/25486 A1 (RANA), WO 02/08337 A1 (ALMSTEAD), and/or WO 02/083353 A1 (RANDO et al.) in view of WANG et al., Nucleic Acids Research Vol. 18, No. 22, HYDE-DREVINYSCHER et al., Chem. & Biol. Vol. 7, No. 1, and Lit et al., Science Vol. 280 (4/1999).

The presently claimed invention is directed to identifying antiproliferative compounds by screening (e.g., high throughput assays) compounds (e.g., library derived) for their ability to inhibit the endouncleodysis of animal IRNA by inhibiting IRNA-IRNA splicing endounclease binding relative to a control.

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Supplemental Box

In case the space in any of the preceding boxes is not sufficient.

Screening assays (e.g., high throughput assays) of single compounds or compound libraries for their ability to disrupt RNA (e.g., RNA) interactions (e.g. including splicing) in order to identify antiproliferative drug candidates is taught by the RANA, ALMSTEAD and/or RANDO reference whose teaching discussed above is hereby incorporated by reference in its entirety.

and/or RANIDO reference whose teaching discussed above is heroby incorporated by reference in its entirety.

The RANA, ALMSTEAD and/or RANDO reference methods differ from the presently claimed invention by failing to explicitly teach the application of its methods to IRNA splicing endonuclease assays that cleave tRNA and tRNA splicing endonuclease.

However, LI et al. teach that the tRNA splicing pathway is analogous in mammals and other organisms (e.g., fung).

In this regard, WANG et al. teach an assay for endonucleolytic tRNA maturation, where inactivated micrococcal nuclease

(reversible inhibito) boand to reliabled pre-RNA physically blocks the sites of endonuclease cleavage and prevents RNA processing activities present in Fraction III of spinach chloroplasts, presumably by substante occlusion or "masking", where formation of an inactive micrococcal nuclease enzyme substante complex precludes withinking to the RNA substrate by a second enzyme.

an inactive micrococcal nuclease enzyme substante complex piecutoses intributor to the 10th Section of "small molecule" compound Additionally, the HYDE-DERUYSCHER et al. reference teaches that high throughput screening of "small molecule" compound libraries (e.g., phage) is ideal for screening "small molecule" enzyme inhibitors for a variety of different enzymes.

Accordingly, it would have been obvious to use fRNA splicing endonuclease assays in the high throughput screening methods of RANA, ALMSTEAD and/or RANDO, because these references specifically suggest screening small molecules libraries for compound which disrupt flatNA interactions, including splicing, and in light of the secondary reference teaching that fRVA's splicing pathway in animals is known and sandogous; and the known teaching of fRNA splicing endonuclease inhibition; with the desirability of using high throughput screening of small molecular libraries for screening express building compromates a drug candidates.